#### CENTER FOR DRUG EVALUATION AND RESEARCH

APPLICATION NUMBER:

20-987

#### FINAL PRINTED LABELING

(pantoprazole sodium)
Delayed-Release Tablets

#### DESCRIPTION

a molecular weight of 432.4. The structural formula is: sesquihydrate, a compound that inhibits gastric acid secretion. Its empirical formula is C<sub>16</sub>H<sub>14</sub>F<sub>2</sub>N<sub>3</sub>NaO<sub>4</sub>S x 1.5 H<sub>2</sub>O, with benzim idazole, so dium 5-(difluoromethoxy)-2-[[(3,4-dimethoxy-2-pyridinyl)methyl] sulfinyl]-1 H-benzim idazole, so dium 5-(difluoromethoxy)-1 H-benzim idazole, so diuThe active ingredient in PROTONIX® (pantoprazole sodium) Delayed-Release Tablets is a substituted

$$OCF_2H$$

in phosphate buffer at pH 7.4, and practically insoluble in n-hexane. weakly basic and acidic properties. Pantoprazole sodium sesquihydrate is freely soluble in water, very slightly soluble Pantoprazole sodium sesquihydrate is a white to off-white crystalline powder and is racemic. Pantoprazole has

220 hours at pH 7.8. decreasing pH. At ambient temperature, the degradation half-life is approximately 2.8 hours at pH 5.0 and approximately The stability of the compound in aqueous solution is pH-dependent. The rate of degradation increases with

acid copolymer NF, polysorbate 80 NF, sodium lauryl sulfate NF, and triethyl citrate NF. hydroxypropyl methylcellulose USP, titanium dioxide USP, yellow iron oxide NF, propylene głycol USP, methacrylic 45.1 mg of pantoprazole sodium sesquihydrate (equivalent to 40 mg pantoprazole) with the following inactive ingredients: anhydrous sodium carbonate NF, mannitol USP, crospovidone NF, povidone USP, calcium stearate NF, PROTONIX is supplied as a delayed-release tablet for oral administration. Each delayed-release tablet contains

Delayed-Release Tablets (pantoprazole sodium)

# CLINICAL PHARMACOLOGY

#### **Pharmacokinetics**

pharmacokinetics are unaltered with multiple daily dosing. Following oral or intravenous administration, the serum administration of pantoprazole to extensive metabolizers, its total clearance is 7.6-14.0 L/h and its apparent volume of pantoprazole is given with food, its t<sub>max</sub> is highly variable and may increase significantly. Following intravenous (t<sub>max</sub>) is 2.4 h and the total area under the plasma concentration versus time curve (AUC) is 4.8 μg·hr/mL. When coated 40 mg pantoprazole tablet, the peak concentration (C<sub>max</sub>) is 2.4 µg/mL, the time to reach the peak concentration In extensive metabolizers (see Metabolism section) with normal liver function receiving an oral dose of the entericconcentration of pantoprazole declines biexponentially with a terminal elimination half-life of approximately one hour in a manner proportional to oral and intravenous doses from 10 mg to 80 mg. Pantoprazole does not accumulate and its leaves the stomach. Peak serum concentration (C<sub>max</sub>) and area under the serum concentration time curve (AUC) increase distribution is 11.0-23.6L. PROTONIX is prepared as an enteric-coated tablet so that absorption of pantoprazole begins only after the tablet

antacids. Administration of pantoprazole with food may delay its absorption up to 2 hours or longer; however, the C<sub>max</sub> absolute bioavailability of approximately 77%. Pantoprazole absorption is not affected by concomitant administration of timing of meals. and the extent of pantoprazole absorption (AUC) are not altered. Thus, pantoprazole may be taken without regard to multiple oral 40-mg doses. Pantoprazole is well absorbed; it undergoes little first-pass metabolism resulting in an The absorption of pantoprazole is rapid, with a C<sub>max</sub> of 2.5 µg/mL that occurs approximately 2.5 hours after single or

fluid. The serum protein binding of pantoprazole is about 98%, primarily to albumin. The apparent volume of distribution of pantoprazole is approximately 11.0-23.6L, distributing mainly in extracellular

is no evidence that any of the pantoprazole metabolites have significant pharmacologic activity. CYP2C19 displays a demethylation, by CYP2C19, with subsequent sulfation; other metabolic pathways include oxidation by CYP3A4. There metabolism is independent of the route of administration (intravenous or oral). The main metabolic pathway is Pantoprazole is extensively metabolized in the liver through the cytochrome P450 (CYP) system. Pantoprazole

(pantoprazole sodium) Delayed-Release Tablets

known genetic polymorphism due to its deficiency in some sub-populations (e.g. 3% of Caucasians and Africanhalf-life values of 3.5 to 10.0 hours, they still have minimal accumulation (≤ 23%) with once daily dosing. Americans and 17-23% of Asians). Although these sub-populations of slow pantoprazole metabolizers have elimination

approximately 71% of the dose was excreted in the urine with 18% excreted in the feces through biliary excretion. There was no renal excretion of unchanged pantoprazole. After a single oral or intravenous dose of <sup>14</sup>C-labeled pantoprazole to healthy, normal metabolizer volunteers.

#### Special Populations

recommended based on age to 76 years of age) after repeated oral administration, compared with younger subjects. No dosage adjustment is Only slight to moderate increases in pantoprazole AUC (43%) and C<sub>max</sub> (26%) were found in elderly volunteers (64

The pharmacokinetics of pantoprazole have not been investigated in patients <18 years of age

clearance values are similar in women and men. No dosage adjustment is needed based on gender (Also see Use in There is a modest increase in pantoprazole AUC and Cmax in women compared to men. However, weight-normalized

#### Renal Impairmen

healthy subjects. No dosage adjustment is necessary in patients with renal impairment or in patients undergoing In patients with severe renal impairment, pharmacokinetic parameters for pantoprazole were similar to those of

#### Hepatic Impairment

patients result in minimal drug accumulation following once daily multiple-dose administration. No dosage adjustment is metabolizers, where no dosage frequency adjustment is warranted. These pharmacokinetic changes in hepatic-impaired by 5- to 7-fold in hepatic-impaired patients, these increases were no greater than those observed in slow CYP2C19 (1.5-fold) relative to healthy subjects. Although serum half-life values increased to 7-9 hours and AUC values increased In patients with mild to moderate hepatic impairment, maximum pantoprazole concentrations increased only slightly

Delayed-Release Tablets (pantoprazole sodium)

other day in these patients. well characterized in patients with severe hepatic impairment. Therefore, the potential for modest drug accumulation needed in patients with mild or moderate hepatic impairment. The pharmacokinetics of pantoprazole have not yet been (≤ 21%) when dosed once daily needs to be weighed against the potential for reduced acid control when dosed every

## **Drug-Drug Interactions**

drug interaction studies with CYP2C19 substrates (diazepam [also a CYP3A4 substrate] and phenytoin [also a CYP3A4 significantly affect the kinetics of other drugs (cisapride, theophylline, diazepam [and its active metabolite, significantly affect the pharmacokinetics of pantoprazole. In vivo studies also suggest that pantoprazole does not altered. It is, therefore, expected that other drugs metabolized by CYPs 2C19, 3A4, 2D6, 2C9 and 1A2 would not theophylline (a CYP1A2 substrate) in healthy subjects, the pharmacokinetics of pantoprazole were not significantly inducer]), nifedipine (a CYP3A4 substrate), metoprolol (a CYP2D6 substrate), diclofenac (a CYP2C9 substrate) and by CYPs 2C19, 3A4, 2C9, 2D6 and IA2. Therefore, it is expected that pantoprazole would not significantly affect the desmethyldiazepam], phenytoin, warfarin, metoprolol, nifedipine, carbamazepine and oral contraceptives) metabolized had no clinically relevant interactions with pantoprazole. they are co-administered with pantoprazole. In other in vivo studies, digoxin, ethanol, glyburide, antipyrine, and caffeine pharmacokinetics of other drugs metabolized by these isozymes. Dosage adjustment of such drugs is not necessary when Pantoprazole is metabolized mainly by CYP2C19 and to minor extents by CYPs 3A4, 2D6 and 2C9. In in vivo drug-

#### **Pharmacodynamics**

#### Mechanism of Action

effect is dose-related and leads to inhibition of both basal and stimulated gastric acid secretion irrespective of the covalent bond to two sites of the (H+,K+)-ATPase enzyme system at the secretory surface of the gastric parietal cell. This stimulus. The binding to the (H<sup>+</sup>,K<sup>+</sup>)-ATPase results in a duration of antisecretory effect that persists longer than 24 hours Pantoprazole is a proton pump inhibitor (PPI) that suppresses the final step in gastric acid production by forming a

#### Antisecretory Activity

occurs after a single dose of oral (20-80 mg) or a single dose of intravenous (20-120 mg) pantoprazole in healthy oral dose of 40 mg pantoprazole, a 51% mean inhibition was achieved by 2.5 hours. With once a day dosing for 7 days volunteers. Pantoprazole given once daily results in increasing inhibition of gastric acid secretion. Following the initial the mean inhibition was increased to 85%. Pantoprazole suppressed acid secretion in excess of 95% in half of the Under maximal acid stimulatory conditions using pentagastrin, a dose-dependent decrease in gastric acid output

of rebound hypersecretion subjects. Acid secretion had returned to normal within a week after the last dose of pantoprazole; there was no evidence

pantoprazole on median pH from one double-blind crossover study are shown below. pantoprazole produced optimal increases in gastric pH which were significantly greater than the 20-mg dose. Doses In a series of dose-response studies pantoprazole, at oral doses ranging from 20 to 120 mg, caused dose-related increases in median basal gastric pH and in the percent of time gastric pH was > 3 and > 4. Treatment with 40 mg of higher than 40 mg (60, 80, 120 mg) did not result in further significant increases in median gastric pH. The effects of

Time	Placebo	20 mg	40 mg	80 mg
8 a.m 8 a.m.	1,3	2.9*	3.8*#	3.9•#
(24 hours)				
8 a.m 10 p.m.	1.6	3.2*	4.4*#	4.8*#
(Daytime)				
10 p.m 8 a.m.	1.2	2.1*	3.0*	2.6*
(Nighttime)				
+ G' 'C' i I'C' i Com - back	3.00	-laaka		
-	Title Committee			

- Significantly different from placebo
- # Significantly different from 20 mg

both one day and one week treatment periods, pantoprazole administered in the morning produced significantly greater increases in median pH during 24 hours than did omeprazole. A double-blind crossover study compared pantoprazole 40 mg with omeprazole 20 mg once daily for 7 days. For

Delayed-Release Tablets (pantoprazole sodium)

## Serum Gastrin Effects

was noted at the 8 week visit with mean increases of 3%, 26%, and 84% for the three pantoprazole dose groups pretreatment values in the 10, 20 and 40 mg treatment groups, respectively. A similar increase in serum gastrin levels up to 8 weeks. At 4 weeks of treatment there was an increase in mean gastrin levels of 7%, 35%, and 72% over (EE) in which 682 patients with gastroesophageal reflux disease (GERD) received 10, 20, or 40 mg of pantoprazole for Fasting serum gastrin levels were assessed in two double-blind studies of the acute healing of erosive esophagitis

generally remained at approximately 2 to 3 times baseline for up to 4 years of periodic follow-up in clinical trials. gastrin level was observed in the initial months of treatment with pantoprazole at doses of 40 mg per day during GERD maintenance studies and 40 mg or higher per day in patients with refractory GERD. Fasting serum gastrin levels In long term studies involving over 800 patients, a 2- to 3-fold mean increase from the pretreatment fasting serum

Following healing of gastric or duodenal ulcers with pantoprazole treatment, elevated gastrin levels return to normal

# Enterochromaffin-Like (ECL) Cell Effects

years, there was a moderate increase in ECL-cell density starting after the first year of use which appeared to plateau In 39 patients treated with oral pantoprazole 40 mg to 240 mg daily (majority receiving 40 mg to 80 mg) for up to 5

tumors .. Gastric NE-cell tumors in rats may result from chronic elevation of serum gastrin levels.. The high density of mg/kg/day resulted in dose-related increases in gastric ECL-cell proliferation and gastric neuroendocrine (NE)-cell concomitant ECL-cell proliferative changes was observed in 1 female rat following 12 months of dosing with administration of pantoprazole at a dose of 0.5 mg/kg/day. In a separate study, a gastric NE-cell tumor without produced by proton pump inhibitors. However, there were no observed elevations in scrum gastrin following the ECL cells in the rat stomach makes this species highly susceptible to the proliferative effects of elevated gastrin levels Impairment of Fertility). pantoprazole at 5 mg/kg/day and a 9-month off-dose recovery.. (See PRECAUTIONS, Carcinogenesis, Mutagenesis, In a nonclinical study in Sprague-Dawley rats, lifetime exposure (24 months) to pantoprazole at doses of 0.5 to 200

(pantoprazole sodium)
Delayed-Release Tablets

#### Other Effects

stimulating hormone, thyronine-binding protein, parathyroid hormone, insulin, glucagon, renin, aldosterone, follicleeffect on the levels of the following hormones: cortisol, testosterone, triiodothyronine (T3), thyroxine (T4), thyroidstimulating hormone, luteinizing hormone, prolactin and growth hormone. function have been detected. In a clinical pharmacology study, pantoprazole 40 mg given once daily for 2 weeks had no No clinically relevant effects of pantoprazole on cardiovascular, respiratory, ophthalmic, or central nervous system

#### Clinical Studies

PROTONIX Delayed-Release Tablets were used in all clinical trials.

# Erosive Esophagitis (EE) Associated with Gastroesophageal Reflux Disease (GERD)

scale). In this study, approximately 25% of enrolled patients had severe EE of grade 3 and 10% had grade 4. The conducted in 603 patients with reflux symptoms and endoscopically diagnosed EE of grade 2 or above (Hetzel-Dent percentages of patients healed (per protocol, n=541) in this study were as follows: A US multicenter double-blind, placebo-controlled study of PROTONIX 10 mg, 20 mg or 40 mg once daily was

	Erosive Esop	hagitis Healing R	Erosive Esophagitis Healing Rates (per protocol)	
		PROTONIX		Placebo
	10 mg QD	20 mg QD	40 mg QD	
Week	(n = 153)	(n = 158)	(n = 162)	(n = 68)
4	45.6%	58.4%+#	75.0%+*	14.3%
∞	66.0%	83.5 %*#	92.6%**	39.7%

<sup>+(</sup>p < 0.001) PROTONIX versus placebo.

# (p<0.05) versus 10 mg PROTONIX

true regardless of H. pylori status for the 20-mg and 40-mg PROTONIX treatment groups. The 40-mg dose of PROTONIX resulted in healing rates significantly greater than those found with either the 20- or 10-mg dose In this study, all PROTONIX treatment groups had significantly greater healing rates than the placebo group. This was

<sup>\*(</sup>p <0.05) versus 10 mg, or 20 mg PROTONIX

nighttime heartburn and the absence of regurgitation starting from the first day of treatment compared with placebo. Patients taking PROTONIX consumed significantly fewer antacid tablets per day than those taking placebo. A significantly greater proportion of patients taking PROTONIX 40 mg experienced complete relief of daytime and

PROTONIX 20 mg and 40 mg once daily was also compared with nizatidine 150 mg twice daily in a US multicenter, double-blind study of 243 patients with reflux symptoms and endoscopically diagnosed EE of grade 2 or above. The percentages of patients healed (per protocol, n=212) were as follows:

41.4%	82.9%	79.2% <sup>†</sup>	∞
22.2%	$64.0\%^{+}$	61.4%	4
(n = 70)	(n = 70)	(n = 72)	Week
150 mg BID	40 mg QD	20 mg QD	٠
Nizatidine	ONIX	PROTONIX	
otocol)	Erosive Esophagitis Healing Rates (per protocol)	osive Esophagitis I	E

<sup>†</sup>(p < 0.001) PROTONIX versus nizatidine.

greater healing rates compared to nizatidine were achieved regardless of the H. pylori status. weeks compared with twice daily treatment with 150 mg of nizatidine. For the 40 mg treatment group, significantly Once daily treatment with PROTONIX 20 or 40 mg resulted in significantly superior rates of healing at both 4 and 8

day than those taking nizatidine nighttime heartburn and regurgitation starting on the first day and of daytime heartburn on the second day compared with those taking nizatidine 150 mg twice daily. Patients taking PROTONIX consumed significantly fewer antacid tablets per A significantly greater proportion of the patients in the PROTONIX treatment groups experienced complete relief of

# INDICATIONS AND USAGE

Short-Term Treatment of Erosive Esophagitis Associated With Gastroesophageal Reflux Disease (GERD)

(pantoprazole sodium)
Delayed-Release Tablets

additional 8 week course of PROTONIX may be considered. symptomatic relief of erosive esophagitis. For those patients who have not healed after 8 weeks of treatment, an PROTONIX Delayed-Release Tablets are indicated for the short-term treatment (up to 8 weeks) in the healing and

(see PRECAUTIONS). The safety and efficacy of PROTONIX for maintenance therapy (e.g., beyond 16 weeks) have not been established.

## CONTRAINDICATIONS

of the formulation. PROTONIX Delayed-Release Tablets are contraindicated in patients with known hypersensitivity to any component

#### **PRECAUTIONS**

#### General

Symptomatic response to therapy with pantoprazole does not preclude the presence of gastric malignancy

animal findings to humans is unknown. The safety and efficacy of PROTONIX for maintenance therapy (e.g., beyond 16 weeks) have not been established. PROTONIX is not indicated for maintenance therapy (see INDICATIONS AND In rodents, pantoprazole is carcinogenic and caused rare types of gastrointestinal tumors. The relevance of these

control when dosed every other day in these patients. pantoprazole has not been well characterized in patients with severe hepatic impairment. Therefore, the potential for modest drug accumulation ( $\leq 21\%$ ) when dosed once daily needs to be weighed against the potential for reduced acid No dosage adjustment is necessary in patients with mild or moderate hepatic impairment. The pharmacokinetics of

(pantoprazole sodium) Delayed-Release Tablets

## **Information for Patients**

affect the absorption of pantoprazole. tablets should be swallowed whole, with or without food in the stomach. Concomitant administration of antacids does not Patients should be cautioned that PROTONIX Delayed-Release Tablets should not be split, crushed or chewed. The

#### Drug Interactions

other drugs metabolized by the cytochrome P450 system, no dosage adjustment is needed with concomitant use of the be necessary. There was also no interaction with concomitantly administered antacids. Therefore, when co-administered with pantoprazole, adjustment of the dosage of pantoprazole or of such drugs may not Clinically relevant interactions of pantoprazole with other drugs with the same metabolic pathways are not expected. glyburide, an oral contraceptive (levonorgestrel/ethinyl estradiol), metoprolol, nifedipine, phenytoin, or warfarin. subsequently undergoes Phase II conjugation. Based on studies evaluating possible interactions of pantoprazole with following drugs: theophylline, cisapride, antipyrine, caffeine, carbamazepine, diazepam, diclofenac, digoxin, ethanol, Pantoprazole is metabolized through the cytochrome P450 system, primarily the CYP2C19 and CYP3A4 isozymes, and

ketoconazole, ampicillin esters, and iron salts). may interfere with absorption of drugs where gastric pH is an important determinant of their bioavailability (e.g. Because of profound and long lasting inhibition of gastric acid secretion, it is theoretically possible that pantoprazole

# Carcinogenesis, Mutagenesis, Impairment of Fertility

gastric fundus at 200 mg/kg/day.. In the liver, treatment at 0.5 to 200 mg/kg/day produced dose-related increases in the malignant neuroendocrine cell tumors in a dose-related manner. In the forestomach, treatment at 50 and 200 mg/kg/day about 0.1 to 40 times the exposure on a body surface basis, of a 50-kg person dosed at 40 mg/day. In the gastric increased incidences of follicular cell adenomas and carcinomas for both male and female rats treatment included an adenocarcinoma of the duodenum at 50 mg/kg/day, and benign polyps and adenocarcinomas of the papillomas and malignant squamous cell carcinomas. Rare gastrointestinal tumors associated with pantoprazole (about 10 and 40 times the recommended human dose on a body surface area basis) produced benign squamous cell fundus, treatment at 0.5 to 200 mg/kg/day produced enterochromaffin-like (ECL) cell hyperplasia and benign and incidences of hepatocellular adenomas and carcinomas. In the thyroid gland, treatment at 200 mg/kg/day produced In a 24-month carcinogenicity study, Sprague-Dawley rats were treated orally with doses of 0.5 to 200 mg/kg/day,

(pantoprazole sodium)
Delayed-Release Tablets

rats exposed to pantoprazole in 6-month and 12-month toxicity studies. Sporadic occurrences of hepatocellular adenomas and a hepatocellular carcinoma were observed in Sprague-Dawley

approximately 1 to 10 times the recommended human dose based on body surface area. In the gastric fundus, treatment at tumors. Dose selection for this study may not have been adequate to comprehensively evaluate the carcinogenic potential 5 to 50 mg/kg/day produced enterochromaffin-like (ECL) cell hyperplasia and benign and malignant neuroendocrine cell In a 24-month carcinogenicity study, Fischer 344 rats were treated orally with doses of 5 to 50 mg/kg/day,

mg/kg/day also produced gastric fundic ECL cell hyperplasia. times the recommended human dose based on body surface area. In the liver, treatment at 150 mg/kg/day produced increased incidences of combined hepatocellular adenomas and carcinomas in female mice. Treatment at 5 to 150 In a 24-month carcinogenicity study, B6C3F1 mice were treated orally with doses of 5 to 150 mg/kg/day, 0.5 to

assay for mutagenic effects. Equivocal results were observed in the in vivo rat liver DNA covalent binding assay. micronucleus tests for clastogenic effects, and in the in vitro Chinese hamster ovarian cell/HGPRT forward mutation mutation test with mouse lymphoma L5178Y cells, and the in vivo rat bone marrow cell chromosomal aberration assay. with rat hepatocytes, the in vitro AS52/GPT mammalian cell-forward gene mutation assay, the in vitro thymidine kinase Pantoprazole was negative in the in vitro Ames mutation assay, , the in vitro unscheduled DNA synthesis (UDS) assay Pantoprazole was positive in the *in vitro* human lymphocyte chromosomal aberration assays, in one of two mouse

surface area) and 450 mg/kg/day in female rats (88 times the recommended human dose based on body surface area) was found to have no effect on fertility and reproductive performance. Pantoprazole at oral doses up to 500 mg/kg/day in male rats (98 times the recommended human dose based on body

### Pregnancy Teratogenic Effects Pregnancy Category B

dose based on body surface area) and rabbits at oral doses up to 40 mg/kg/day (16 times the recommended human dose are not always predictive of human response, this drug should be used during pregnancy only if clearly needed There are, however, no adequate and well-controlled studies in pregnant women. Because animal reproduction studies based on body surface area) and have revealed no evidence of impaired fertility or harm to the fetus due to pantoprazole Teratology studies have been performed in rats at oral doses up to 450 mg/kg/day (88 times the recommended human

(pantoprazole sodium)
Delayed-Release Tablets

#### **Nursing Mothers**

should be made whether to discontinue nursing or to discontinue the drug, taking into account the benefit of the drug to the infants. Based on the potential for tumorigenicity shown for pantoprazole in rodent carcinogenicity studies, a decision human milk. Many drugs which are excreted in human milk have a potential for serious adverse reactions in nursing Pantoprazole and its metabolites are excreted in the milk of rats. It is not known whether pantoprazole is excreted

#### **Pediatric Use**

Safety and effectiveness in pediatric patients have not been established

#### Use in Women

those found in men. The incidence rates of adverse events were also similar between men and women Erosive esophagitis healing rates in the 221 women treated with pantoprazole in US clinical trials were similar to

#### Use in Elderly

age. The healing rates of the 25 patients at least 75 years old were 80% for those treated with 10 mg of pantoprazole and abnormalities in patients aged 65 years and older were similar to those associated with patients younger than 65 years of similar to that of patients younger than 65 years of age. trials were similar to those found in patients under the age of 65. The incidence rates of adverse events and laboratory 100% for those patients treated with either 20 or 40 mg. In addition, the safety profile in patients 65 years and older was Erosive esophagitis healing rates in the 107 elderly patients (≥65 years old) treated with pantoprazole in US clinical

## ADVERSE REACTIONS

dosages and duration of treatment. In general, pantoprazole has been well tolerated in both short-term and long-term Worldwide, more than 11,100 patients have been treated with pantoprazole in clinical trials involving various

possibly, probably or definitely related to drug occurred in 1% or more in the individual studies of GERD patients on dose-related effects on the incidence of adverse events. The following adverse events considered by investigators to be therapy with PROTONIX. In two US controlled clinical trials involving PROTONIX 10-, 20-, or 40-mg doses for up to 8 weeks, there were no

#### Delayed-Release Tablets (pantoprazole sodium)

Most Frequent Adverse Events Reported as Drug Related in Short-term Domestic Trials ----- % Incidence -----

	Study 300-US	)0-US	Study 301-US	01-US
	PROTONIX	Placebo	PROTONIX	Nizatidine
Study Event	(n = 521)	(n = 82)	(n = 161)	(n = 82)
Headache	6	6	9	13
Diarrhea	4	_	6	6
Flatulence	2	2	4	0
Abdominal pain	-	2	4	4
Rash	<u>^</u>	0	2	0
Eructation	_	_	0	0
Insomnia	<u>^</u>	2		_
Hyperglycemia	_	0	<u>-</u>	0

and urinary tract infection. bronchitis, cough increased, dyspnea, pharyngitis, rhinitis, sinusitis, upper respiratory tract infection, urinary frequency, vomiting, hyperlipemia, liver function tests abnormal, SGPT increased, arthralgia, anxiety, dizziness, hypertonia, infection, pain, migraine, constipation, dyspepsia, gastroenteritis, gastrointestinal disorder, nausea, rectal disorder, occurred at a rate of ≥ 1% in PROTONIX-treated patients: asthenia, back pain, chest pain, neck pain, flu syndrome, In addition, in these short-term domestic trials, the following treatment-emergent events, regardless of causality,

adverse events were reported to occur in 1% or more of 2805 GERD patients receiving pantoprazole for up to 8 weeks In international short-term double-blind or open-label, clinical trials involving 20- to 80 mg per day, the following

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Adverse Events in GERD Patients in Short-term International Trials
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Abdominal Pain	Diarrhea	Headache	Study Event				Auverse
_	2	2	(N=2805)	Total	Pantoprazole		Adverse Evens in CERD Fatients in Silon-term international infats
_	2	3	(N=594)	300 mg	Ranitidine	% Inc	cous ill short-ter
^	2	2	(N=474)	20 mg	Omeprazole	cidence	II THERMANIAL TI
<u>^</u>	^	1	(N=239)	40 mg	Famotidine	1	Ials
	Abdominal Pain 1 <1 <1	Diarrhea 2 2 <1  Abdominal Pain 1 1 <1 <1	Headache       2       3       2       1         Diarrhea       2       2       2       <1	(N=2805) (N=594) (N=474) 2 3 2 2 2 2 2 1 1 <1	Total 300 mg 20 mg (N=2805) (N=594) (N=474) 2 3 2 2 1 1 <1	Pantoprazole Ranitidine Omeprazole Total 300 mg 20 mg (N=2805) (N=594) (N=474)  2 3 2 2 2 2 1 1 < 1	Pantoprazole Ranitidine Omeprazole Total 300 mg 20 mg (N=2805) (N=594) (N=474)  2 3 2 1 1 < 1

pantoprazole was unclear. domestic or international trials are shown below within each body system. In most instances the relationship to Additional adverse experiences occurring in <1% of GERD patients based on pooled results from either short-term

laboratory test abnormal, malaise, moniliasis, neoplasm, non-specified drug reaction BODY AS A WHOLE: abscess, allergic reaction, chills, cyst, face edema, fever, generalized edema, heat stroke, hernia

retinal vascular disorder, syncope, tachycardia, thrombophlebitis, thrombosis, vasodilatation. heart failure, electrocardiogram abnormal, hemorrhage, hypertension, hypotension, myocardial ischemia, palpitation. CARDIOVASCULAR SYSTEM: angina pectoris, arrhythmia, cardiovascular disorder, chest pain substernal, congestive

esophageal hemorrhage, esophagitis, gastrointestinal carcinoma, gastrointestinal hemorrhage, gastrointestinal moniliasis, abscess, periodontitis, rectal hemorrhage, stomach ulcer, stomatitis, stools abnormal, tongue discoloration, ulcerative gingivitis, glossitis, halitosis, hematemesis, increased appetite, melena, mouth ulceration, oral moniliasis, periodontal DIGESTIVE SYSTEM: anorexia, aphthous stomatitis, cardiospasm, colitis, dry mouth, duodenitis, dysphagia, enteritis,

ENDOCRINE SYSTEM: diabetes mellitus, glycosuria, goiter.

alkaline phosphatase increased, gamma glutamyl transpeptidase increased, SGOT increased HEPATO-BILIARY SYSTEM: biliary pain, bilirubinemia, cholecystitis, cholelithiasis, cholestatic jaundice, hepatitis,

(pantoprazole sodium)
Delayed-Release Tablets

leukocytosis, leukopenia, thrombocytopenia. HEMIC AND LYMPHATIC SYSTEM: anemia, ecchymosis, eosinophilia, hypochromic anemia, iron deficiency anemia,

METABOLIC AND NUTRITIONAL: dehydration, edema, gout, peripheral edema, thirst, weight gain, weight loss

rigidity, myalgia, tenosynovitis MUSCULOSKELETAL SYSTEM: arthritis, arthrosis, bone disorder, bone pain, bursitis, joint disorder, leg cramps, neck

decreased, sleep disorder, somnolence, thinking abnormal, tremor, vertigo. hallucinations, hyperkinesia, hypesthesia, libido decreased, nervousness, neuralgia, neuritis, paresthesia, reflexes NERVOUS SYSTEM: abnormal dreams, confusion, convulsion, depression, dry mouth, dysarthria, emotional lability,

RESPIRATORY SYSTEM: asthma, epistaxis, hiccup, laryngitis, lung disorder, pneumonia, voice alteration

simplex, herpes zoster, lichenoid dermatitis, maculopapular rash, pain, pruritus, skin disorder, skin ulcer, sweating, SKIN AND APPENDAGES: acne, alopecia, contact dermatitis, dry skin, eczema, fungal dermatitis, hemorrhage, herpes

glaucoma, otitis externa, taste perversion, tinnitus. SPECIAL SENSES: abnormal vision, amblyopia, cataract specified, deafness, diplopia, ear pain, extraocular palsy,

urethritis, urinary tract disorder, urination impaired, vaginitis. impotence, kidney calculus, kidney pain, nocturia, prostatic disorder, pyelonephritis, scrotal edema, urethral pain, UROGENITAL SYSTEM: albuminuria, balanitis, breast pain, cystitis, dysmenorrhea, dysuria, epididymitis, hematuria,

## **Postmarketing Reports**

include anaphylaxis; angioedema (Quincke's edema); anterior ischemic optic neuropathy; severe dematologic reactions, pancreatitis. including erythema multiforme, Stevens-Johnson syndrome, and toxic epidermal necrolysis (TEN, some fatal); and There have been spontaneous reports of adverse events with the post-marketing use of pantoprazole. These reports

(pantoprazole sodium)
Delayed-Release Tablets

nausea, and tinnitus In addition, also observed have been jaundice, confusion, hypokinesia, speech disorder, increased salivation, vertigo,

#### Laboratory Values

hypercholesterolemia, and hyperuricemia alternative explanation for a laboratory value change, such as intercurrent illness, the elevations tended to be mild and sporadic. The following changes in laboratory parameters were reported as adverse events: creatininc increased three times the upper limit of normal at the final treatment visit. Except in those patients where there was a clear In two US controlled trials, 0.4 % of the patients on 40 mg pantoprazole experienced SGPT elevations of greater than

#### OVERDOSAGE

unknown doses of chloroquine and zopiclone which were also taken since two other reported cases of pantoprazole subjects and have been well tolerated. well tolerated. Doses of up to 240 mg per day, given intravenously for seven days, have been administered to healthy a flexible dosing study of refractory peptic ulcer disease received a dose of 320 mg per day for 3 months; treatment was overdosage involved similar amounts of pantoprazole (400 and 600 mg) with no adverse effects observed. One patient in overdosage of pantoprazole (560 mg) has been received; however, the death was more reasonably attributed to the Some reports of overdosage with pantoprazole have been received. A spontaneous report of a suicide involving an

Pantoprazole is not removed by hemodialysis.

segregation, absence of ear reflex, and tremor. respectively. The symptoms of acute toxicity were hypoactivity, ataxia, hunched sitting, limb-splay, lateral position, Single oral doses of pantoprazole at 709 mg/kg, 798 mg/kg and 887 mg/kg were lethal to mice, rats and dogs,

# DOSAGE AND ADMINISTRATION

# Treatment of Erosive Esophagitis

The recommended adult oral dose is 40 mg given once daily for up to 8 weeks. For those patients who have not healed after 8 weeks of treatment, an additional 8-week course of PROTONIX may be considered. (See INDICATIONS

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once daily needs to be weighed against the potential for reduced acid control when dosed every other day in these patients with severe hepatic impairment. Therefore, the potential for modest drug accumulation ( $\leq 21\%$ ) when dosed mild or moderate hepatic impairment. The pharmacokinetics of pantoprazole have not yet been well characterized in No dosage adjustment is necessary in patients undergoing hemodialysis. No dosage adjustment is needed in patients with No dosage adjustment is necessary in patients with mild, moderate or severe renal insufficiency or in elderly patients.

Concomitant administration of antacids does not affect the absorption of PROTONIX PROTONIX Delayed-Release Tablets should be swallowed whole, with or without food in the stomach

Patients should be cautioned that PROTONIX Delayed-Release Tablets should not be split, chewed or crushed

#### HOW SUPPLIED

ink) on one side. PROTONIX is supplied as 40 mg yellow oval biconvex delayed-release tablets imprinted with PROTONIX (brown

They are available as follows:

NDC 0008-0841-81 bottles of 90

Storage

[See USP Controlled Room Temperature]. Store PROTONIX Delayed-Release Tablets at 20°-25°C (68°-77°F); excursion permitted to 15°-30°C (59°-86°F).

Rx only

US Patent No. 4,758,579

(pantoprazole sodium)
Delayed-Release Tablets

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